Please enter the following amended claim set and consider the remarks herein.

# IN THE CLAIMS

#### Please cancel claims 1-14.

# Please add new claims 15-38 as follows:

- 15.(new) A method of genotyping an allele of a target nucleic acid sequence comprising:
  - a) providing a target nucleic acid sequence comprising a first domain and a second domain, wherein said first and said second domains are separated by a detection position;
  - b) forming a first hybridization complex by:
    - i) hybridizing a first primer comprising an adapter sequence to said first domain, immediately adjacent to and 3' of said detection position;
    - ii) hybridizing a second primer to said second domain, immediately adjacent to and 5' of said detection position;
  - c) contacting said first hybridization complex with dNTPs and a first enzyme to form a modified hybridization complex;
  - d) contacting said modified hybridization complex with a second enzyme to form a ligated probe; and
  - e) detecting the presence of said ligated probe.
- 16.(new) The method according to claim 15, wherein said first enzyme is a polymerase.
- 17.(new) The method according to claim 15, wherein said second enzyme is a ligase.
- 18.( new) The method according to claim 15, wherein said dNTPs comprise a label.
- 19.(new) The method according to claim 18 wherein said label is a fluorescent label.
- 20.(new) The method according to claim 15 further comprising:
  - f) contacting said ligated probe with an array comprising:
    - i) a substrate with a surface comprising discrete sites; and
    - ii) a population of microspheres comprising at least a first subpopulation

comprising a first capture probe, such that said first capture probe and said ligated probe comprising said adapter sequence to form a second hybridization complex; wherein said microspheres are distributed on said surface; and

- g) detecting the presence of said second hybridization complex.
- 21.(new) The method according to claim 20, wherein said substrate is a fiber optic bundle.
- 22.(new) The method according to claim 20, wherein said substrate is selected form the group consisting of glass and plastic.
- 23.(new) The method according to claim 20, wherein said discrete sites comprise wells.
- 24.(new) The method according to claim 15 wherein said second primer further comprises an adapter sequence.
- 25.(new) The method according to claim 15 further comprising:
- f) contacting said ligated probe with an ordered array, wherein said ordered array comprises capture probes.
- 26.(new) The method according to claim 15 further comprising:
- f) contacting said ligated probe with a population of microspheres comprising at least a first subpopulation comprising a first capture probe, such that said first capture probe and said ligated probe comprising said adapter sequence form a second hybridization complex.

## 27.(new) A method comprising:

- a. providing a hybridization complex comprising:
- i) a target nucleic acid comprising a first and a second domain separated by a detection position;
- ii) a first probe hybridized to said first domain, wherein said first probe further comprises an adapter sequence that is not complementary to said target sequence; and
  - iii) a second probe hybridized to said second domain;
- b. contacting said hybridization complex with nucleotides and an extension enzyme such that said first probe is extended to form an extended first probe comprising a nucleotide that is complementary to said detection position, whereby a modified hybridization complex is formed;

- c. contacting said modified hybridization complex with a ligase, whereby said extended first probe and said second probe are ligated forming a ligated probe; and
  - d) detecting said ligated probe.
- 28.(new) The method according to claim 27, wherein said detecting comprises:

contacting said ligated probe with a first immobilized capture probe, whereby said adapter hybridizes with said capture probe.

- 29.(new) The method according to claim 28, wherein said capture probe is immobilized on an ordered array.
- 30.(new) The method according to claim 28, wherein said capture probe is immobilized on a first population of microspheres.
- 31.(new) The method according to claim 28, wherein said capture probe is immobilized on a first population of microspheres, wherein said microspheres are randomly distributed on a substrate.
- 32.(new) The method according to claim 31, wherein said substrate is a fiber optic bundle.
- 33.(new) The method according to claim 27, wherein said substrate is selected from the group consisting of glass and plastic.
- 34.(new) The method according to claim 27 wherein said nucleotides are dNTPS
- 35.(new) The method according to claim 34, wherein said dNTPs are labeled
- 36.(new) The method according to claim 27, wherein said second probe comprises a label.
- 37.(new) The method according to claim 27, wherein said extension enzyme is a DNA polymerase.

#### IN THE SPECIFICATION

Please amend the specification as follows:

On page one of the specification, beginning at line 3, under the title, applicants respectfully request that the claim of priority be changed to read the following: